Kinetic of thermal degradation of alginate-zeolite composites

ABSTRACT

In this work, natural and Fe(III)-modified zeolite were tested as additives for alginate in order to improve its thermal and pyrolytic properties. For that purpose the natural zeolite (NZA) and Fe(III)-modified zeolite coated with alginate (FeA) were used. Thermal degradation of the samples was investigated by using thermal analytic techniques, thermogravimetric (TG) and differential thermal analysis (DTA). The samples were heated from 25°C to 1000°C at different heating rates (from 5 to 20°C). From experimental results, thermal degradation of the sodium alginate could be clearly separated into several stages based on the weight loss and the weight loss rate. The kinetic parameters (activation energy and pre-exponential factor) for the main degradation phase (from 180 to 500°C) thus were estimated by using the pseudo first-order assumption. The results showed that used zeolites affected the thermal properties of the alginate and reduce activation energy required for pyrolysis from 87.30 kJ/mol for pure alginate to 72.75 kJ/mol for NZA and 63.35 kJ/mol for FeA. These pyrolysis investigations can be useful for better understanding the thermal stability of these alginate biocomposites and provide necessary information for thermally processing wastes from these biodegradable materials.

Keywords: Na-alginate, natural zeolite, Fe(III)-modified zeolite, composite, thermogravimetric and differential thermal analysis, thermal degradation kinetic.

1. INTRODUCTION

Nowadays plastics are very often used due to their durability, versatility as well as low price. Using of plastic products give a lot of benefits to humans and modern society in terms of economic activity, jobs and quality of life. Even more, plastics can reduce energy consumption and greenhouse gas emissions in many circumstances, even in some packaging applications when compared to the alternatives. On the other side, plastic is usually non-biodegradable and therefore can remain as waste in the environment for a very long time, it may pose risks to human health as well as the environment and also it can be difficult to reuse and/or recycle in practice [1]. From that point, developing of the new recycle techniques or degradable polymers is economic and environmental necessity. One of possible ways for solving the environmental problems caused by the massive use of synthetic polymer materials and their wastes is developing materials based on biopolymers as well as extending their use [2].

Alginate is a naturally occurring polysaccharide obtained from marine brown algae, comprising linear chains of (1,4)-b-D-mannuronic acid and (1,3)-a-L-guluronic acid. Sodium alginate is a poly-electrolyte and it is biodegradable and non-toxic polymer, with gelation and high water absorption properties. It has rigid molecular structure, and has been extensively studied on many potential applications in medicine, pharmacy, in food industry or for environmental protection in terms of removal of heavy metals from contaminated waters [3, 4]. Also, alginate possesses good film forming ability. Hence, it could be used for food packing as a replacement of conventional synthetic polymers.

One of possible ways for recycling and for avoiding huge accumulation of plastic wastes obtained from packing industry is thermal pyrolysis which may provide a sustainable way of recycling organic wastes. This technique offers the possibility of transforming packing wasters into fuel oils for the petrochemical industries or decomposing these wastes for thermal power generation [5,6].

However, the low thermal conductivity of plastics as well as the endothermic character of the
decomposition reactions involves high power consumption in the pyrolysis process. Using of the additives may permit decrease of the temperature of the process and also could provide a narrower product distribution, which allows control of the type of products obtained with a lower production cost [7]. According to the literature [8], zeolites may act as catalysts in the cracking of polymers, making their transformation into fuel products easier. On the other side, in our previous paper [9] composites of the zeolite-clinoptilolite and alginate (natural (NZA) or Fe(III)-modified zeolite (FeA)) were tested as adsorbents for removal of lead ions from contaminated water solutions. Results obtained for the highest investigated initial concentration of lead (15.1mmol/dm$^3$) showed good adsorption capacities and maximum adsorbed amounts of lead were 0.50 and 0.66mmol/dm$^3$ for NZA and FeA, respectively. Also during adsorption process there was no water turbidity what is beside good adsorption properties very important for experiments in column and application under real conditions. Thus, in this work, thermal degradation of the samples used for adsorption experiments (NZA and FeA) was performed and results were compared with those obtained for pure sodium alginate. The aim was to investigate the influence of the natural and Fe(III)-modified zeolite-clinoptilolite on thermal degradation and pyrolytic properties of the alginate, i.e. indication whether natural and Fe(III)-modified zeolite can be used as additives for thermal decomposition of the alginate.

2. EXPERIMENTAL

Chemicals used in this study were of high purity and supplied from Sigma Aldrich Germany. The natural zeolite-clinoptilolite from Serbia with a particle size <0.043mm (NZ) was used as a starting material. The Fe(III)-modified zeolite (FeZ) was obtained combining the method for the goethite preparation and that for the preparation of Fe-coated zeolite [10, 11], and detailed procedure is given elsewhere [12].

Natural and Fe(III)-modified zeolite-alginate composites were prepared using a method described by Yuan and Viraraghavan [13]. A 2g of Na-alginate was dissolved in 100cm$^3$ of distilled water and then agitated for 24h. Then, 10g of the NZ or FeZ was blended with Na-alginate solution for 2h. Mixed solution was dropped into 0.1mol/dm$^3$ of CaCl$_2$ solution. The beads were obtained and hardened by placing them in 2% solution of CaCl$_2$ for 24h. Then, beads were 5 times washed with distilled water by agitation at 100rpm for 30min. At the end, beads were dried at room temperature. Finally, the NZA (natural zeolite-alginate) and FeA (Fe(III)-modified-alginate beads) were obtained.

Thermal behaviour of the composite samples was followed by differential thermal analysis (DTA) on a Netzsch STA 409 EP. To compare results, DTA and thermogravimetric analysis (TGA) were performed on Na-alginate. Samples were heated from 25 to 1000$^\circ$C in an air atmosphere at heating rates of 5, 10, 15 and 20$^\circ$C/min. The samples were kept in a desiccator at a relative humidity of 23%, prior to analyses.

3. RESULTS AND DISCUSSION

Figure 1 shows the DTA/TGA curves of the pure Na-alginate heated at 5$^\circ$C/min up to 1000$^\circ$C.

![Figure 1 - TGA and DTA curve of the pure Na-alginate at heating rate of 5$^\circ$C/min.](image-url)
As can be seen, DTA graph of Na-alginate shows sixth distinct peaks. An endothermic DTA peak at 113°C is due to dehydration, and due to this process at TGA curve a mass loss of about 15% was noted in temperature interval 25-180°C. The major degradation of Na-alginate occurred in second temperature interval from 180 to 500°C (weight loss of about 50%) where loss of volatile components, rupture of chains and fragmentation of sodium alginate occurs. This process is followed with two exothermic peaks at DTA curve with maximums at 240 and 348°C. Decomposition of Na-alginate and its fragmentation into monomers occurred in third temperature interval form 500 to 700°C and this process was followed with exothermic peaks with maximums at 576 and 598°C and with the weight loss of about 17%. These processes are followed by fragments and monomer units conversion into carbonate as a by-product. Finally due to oxidation of the carbonate, endothermic peak with minimum at 817°C and mass loss of ~6% was occurred in the temperature interval form 700 to 1000°C [16, 17].

From Figure 1 it is clear that processes occurred in the temperature interval from 180 to 700°C are exothermic, indicating that energy released from burning or forming new chemical bonds was greater than the energy absorbed for bond scission during decomposition. That excess between released and activation energy may be collected and used for other energy applications. In order to investigate influence of the zeolites on activation energy for thermal degradation of the alginate, thermal decomposition kinetic was followed for NZA, FeA and Na-alginate and results obtained with differential thermal analysis (DTA) at four different heating rates (5, 10, 15 and 20°C/min) are shown at Figure 2.
Since the main weight loss (the highest part of alginate burned) was observed in temperature interval from 180 to 500°C (~50%), kinetic calculations were performed for most clearly expressed peak at DTA curve in that temperature range (Figure 1, DTA peak at 348°C).

As it can be seen from Figure 2 for all three samples as the heating rate increased the peak of interest became wider and decomposition temperature shifted to the higher values (in DTA diagrams, for Na-alginate from 348 to 395°C, NZA from 327 to 384°C and for FeA from 320 to 364°C at heating rate 5°C and 20°C, respectively). It is very important to notice for composite samples, for the same heating rate, the peak temperatures were lower in comparison with Na-alginate, and also, slightly lower temperatures were observed for FeA than for NZA.

Kinetic parameters, activation energy \( (E_a) \) and pre-exponential factor \( (A) \) for the thermal decomposition of the samples are determined from data obtained by DTA analysis using the Kissinger method [18]. According to Kissinger method thermal decomposition of sample can be described with following first order kinetic equation:

\[
\frac{\partial x}{\partial t} = k_T (1 - x)
\]  

(1)

where, \( x \) is the fraction of material decomposed. The magnitude of the rate constant, \( k_T \), can be determined by the temperature and is given by the Arrhenius equation:

\[
k_T = A e^{-\frac{E_a}{RT}}
\]  

(2)

where \( R \) is gas constant and \( T \) is the Kelvin temperature. Following this method, the relationship between the heating rate \( (\beta) \) and the peak temperature \( (T) \) can be plotted. Obtained results given in Figure 3 showed that the \( \ln(\beta/T_p^2) \) against \( 1/T \) are straight lines for all samples \( (R^2>0.95) \), indicating that the kinetic mechanism of thermal decomposition of these compounds followed the first order kinetics. The activation energy was determined from slope of the lines (the slope is equal to \( -E_a/R \)), while the pre-exponential factor \( \ln(A) \) was calculated from the following equation [19]:

\[
A = \beta \left( \frac{E_a}{RT_p^2} \right) \exp \left( \frac{E_a}{RT_p} \right)
\]  

(3)

and results are listed in Table 1.

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<th>Table 1 - Results obtained by Kissinger method</th>
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<td>( \ln(A) ), min(^{-1} )</td>
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<td>Na-alginate</td>
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Figure 2 - Influence of the heating rate on the DTA curve for: a) Na-alginate, b) NZA c) FeA.
As can be seen from Table 1 the activation energy required for thermal degradation of alginate decreased for composite samples in comparing to the starting Na-alginate. However, lower value of the $E_a$ was obtained for FeA than for NZA. The $E_a$ value decreased from 87.30 kJ/mol for Na-alginate to 72.75 kJ/mol for NZA and to 63.35 kJ/mol for FeA sample. From that point it can be indicated that both samples, NZA and especially FeA, could be used as additives and may render the materials processing temperature and burning activation energy decrease and possibly allow lowering of the thermal treatment cost.

4. CONCLUSION

In this paper thermal degradation of pure alginate and its composites with natural (NZA) and Fe(III)-modified zeolite (FeA) was investigated. From results of thermogravimetric (TG) and differential thermal analysis (DTA) of the pure alginate it was shown that the highest part of alginate burned in temperature interval from 180 to 500°C (~50%) where loss of volatile components, rupture of chains and fragmentation of sodium alginate occurs.

Kinetic calculations were performed for most clearly expressed peak at DTA curve in that temperature range (DTA peak at 348°C). Obtained results showed that decomposition shifts to higher temperature as the heating rate increases and also to the lower temperatures for composite samples in comparison with starting alginate sample.

Kinetic analysis shows that the main thermal degradation follows the first order reaction mechanism. Kissinger method was used for determination of the kinetic parameters (pre-exponential factor and activation energy) from DTA data. Results showed that activation energy for the thermal decomposition of the alginate decreases after addition of the zeolites. Lower values of the activation energy obtained for composite samples indicated that both sample (natural and due to lower value of activation energy, especially Fe(III)-modified zeolite) may be used as additives for thermal degradation of the alginate.

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5. REFERENCES


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IZVOD

KINETIKA TERMINSKOG RAZLAGANJA KOMPOZITA ALGINATA I ZEOLITA

U ovom radu ispitavana je upotreba prirodnog i Fe(III)-modifikovanog zeolita kao aditiva alginate u cilju poboljšanja njegovih termijskih i pirolitičkih osobina. Za tu namenu korišćeni su uzorci pridruženih alginatama (FeA). Sagorevanje uzoraka pruženo je kemijskom analitickim tehnikama, diferencijalnom termijskom analizom (DTA) i termogravimetrijskom analizom (TGA). Uzorci su zagrevani u temperaturskom intervalu od 25 do 727,75 kJ/mol za PZA i 1000 kJ/mol za FeA kao i da smanjuju aktivacionu energiju potrebnu za njegovu pirolizu sa 87,30 kJ/mol za čist alginat do 500 kJ/mol za PZA i FeA. Na osnovu eksperimentalnih rezultata, merenjem gubitka mase, termijsko razlaganje natrijum alginata se jasno može podeliti u nekoliko koraka. Na osnovu kinetičkog modela pseudo-prvog reda određeni su kinetički parametri (aktivacija energija i pred eksponencijalni faktor) koji se odnose na temperaturski interval od 180 do 500°C. Dobijeni rezultati su pokazali da upotrebljeni zeoliti utiču na termijske osobine alginate kao i da smanjuju aktivacionu energiju potrebnu za njegovu pirolizu sa 87,30 kJ/mol za čist alginit do 72,75 kJ/mol za PZA i 63,35 kJ/mol za FeA. Ova ispitivanja mogu biti korisna za bolje razumevanje termijske stabilnosti alginitnih biokompozita i mogu dati informacije važne za termijske prozese biogradujućih materijala.

Ključne reči: Na-alginat, prirodni zeolit, Fe(III)-modifikovani zeolit, kompozit, termogravimetrijska i diferencijalna termijska analiza, kinetika termijske degradacije.

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